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Total No. of Pages : 01

Total No. of Questions : 06

M.Pharmacy (Pharmaceutical Analysis) (2017 &amp; Onwards) (Sem.-1)

**ADVANCED PHARMACEUTICAL ANALYSIS**

Subject Code : MPA-102T

M.Code : 74694

Time : 3 Hrs.

Max. Marks: 75

**INSTRUCTIONS TO CANDIDATES :**

1. Attempt any FIVE questions out of SIX questions.
2. Each question carries FIFTEEN marks.

**1. Explain in brief :**

- |  |     |
|--|-----|
| a. Residual impurities   | 3   |
| b. Application of immunoassay  | 3   |
| c. Production of antibodies  | 3   |
| d. Container orientation in stability testing  | 3   |
| e. Residual solvents   | 3   |
| 2. Give a detailed account on principle, classification, limits and analytical procedure for reporting of residual impurities. | 15  |
| 3. a. Discuss in detail the classification and control of elemental impurities in Pharmaceutical products.                     | 10  |
| b. Give methods of C, H, N and S analysis with principle.  | 5   |
| 4. a. Describe the ICH stability guidelines for biological products. Short note on Stability zones.                            | 5,5 |
| b. Briefly describe the separation techniques in Immunoassays.   | 5   |
| 5. Write in detail about :   |     |
| a. The regulatory requirements for HPTLC finger printing in stability testing of Phytopharmaceuticals.                         | 10  |
| b. Biological tests and assay of Oxytocin  | 5   |
| 6. Write short note on :   |     |
| a. PCR and its application.  | 10  |
| b. Principle involved, procedure and application of Radioimmunoassay   | 5   |

**NOTE : Disclosure of Identity by writing Mobile No. or Making of passing request on any page of Answer Sheet will lead to UMC against the Student.**

